

Dentz S.N. 081595158

=> d his study); PREP (Preparation); USES (Uses)

7-7-1996

(FILE 'HOME' ENTERED AT 11:01:20 ON 08 JUL 96)

FILE 'REGISTRY' ENTERED AT 11:01:58 ON 08 JUL 96

L1		STRU
L2	0	S L1
L3	0	S L1 FUL
L4		STRU 1
L5	0	S L4
L6		STRU L1
L7	0	S L6
L8		STRU L7
L9	2	S L8
L10	67	S L8 FUL
L11		STRU L8
L12	39	SEARCH L11 SUB=L10 FUL

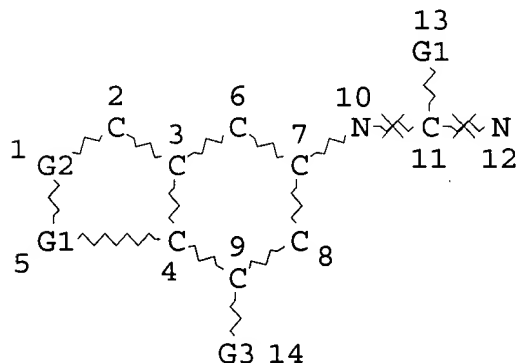
FILE 'CA' ENTERED AT 11:31:22 ON 08 JUL 96

L13 14 S L12

=> d 18

'L8' HAS NO ANSWERS

L8 STR



VAR G1=O/S

REP G2=(1-3) CH

VAR G3=H/AK

NODE ATTRIBUTES:

NSPEC TS BC AT 12

DEFUALT ML-EVEL: IS ATOM

DEFASSET REEVEES IS ATOM
DEFAULT ECL-EVEL IS LIMITED

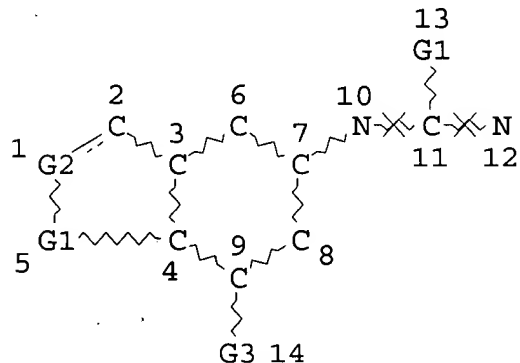
GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED
NUMBER OF NODES IS 14

STEREO ATTRIBUTES: NONE

=> d 19
NO ANSWERS DISPLAYED.
THE ANSWER SET WAS CREATED IN FILE 'REGISTRY'.
USE THE FILE COMMAND TO CHANGE TO THE CORRECT FILE.
You have entered a file that is not in the current file
environment.
Enter "DISPLAY HISTORY" to see a list of the files in the
current
environment.

=> d 111
'L11' HAS NO ANSWERS
L11 STR



VAR G1=O/S
REP G2=(1-3) CH
VAR G3=H/AK
NODE ATTRIBUTES:
NSPEC IS RC AT 12
DEFAULT MLEVEL IS ATOM
DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:
RING(S) ARE ISOLATED OR EMBEDDED
NUMBER OF NODES IS 14

STEREO ATTRIBUTES: NONE

=> d 112 reg can
NO ANSWERS DISPLAYED.
THE ANSWER SET WAS CREATED IN FILE 'REGISTRY'.
USE THE FILE COMMAND TO CHANGE TO THE CORRECT FILE.
You have entered a file that is not in the current file
environment.
Enter "DISPLAY HISTORY" to see a list of the files in the
current
environment.

=> fil reg
FILE 'REGISTRY' ENTERED AT 11:36:43 ON 08 JUL 96

USE IS SUBJECT TO THE TERMS OF YOUR CUSTOMER AGREEMENT
COPYRIGHT (C) 1996 American Chemical Society (ACS)

STRUCTURE FILE UPDATES: 6 JUL 96 HIGHEST RN 177929-67-8
DICTIONARY FILE UPDATES: 6 JUL 96 HIGHEST RN 178150-05-5

TSCA INFORMATION NOW CURRENT THROUGH DECEMBER 1995

Please note that search-term pricing does apply when
conducting SmartSELECT searches.

*** YOU HAVE NEW MAIL ***

'.REGISTRY' IS DEFAULT FORMAT FOR 'REGISTRY' FILE

=> d 112 1- reg can

1 RN 174798-58-4 REGISTRY

REFERENCE 1: 124:261061
2 RN 174798-57-3 REGISTRY

REFERENCE 1: 124:261061
3 RN 149866-40-0 REGISTRY

REFERENCE 1: 119:139079
4 RN 149620-94-0 REGISTRY

REFERENCE 1: 119:160327
5 RN 149620-93-9 REGISTRY

REFERENCE 1: 119:160327
6 RN 133077-32-4 REGISTRY

REFERENCE 1: 114:164000
7 RN 130105-31-6 REGISTRY

REFERENCE 1: 113:191969
8 RN 130105-30-5 REGISTRY

REFERENCE 1: 113:191969
9 RN 130104-50-6 REGISTRY

REFERENCE 1: 113:191969
10 RN 125704-84-9 REGISTRY

REFERENCE 1: 112:118858
11 RN 121456-04-0 REGISTRY

REFERENCE 1: 111:39359
12 RN 103082-23-1 REGISTRY

REFERENCE 1: 111:39359

REFERENCE 2: 105:37517

13	RN	103082-22-0	REGISTRY
REFERENCE	1:	111:39359	
REFERENCE	2:	105:37517	
14	RN	103082-21-9	REGISTRY
REFERENCE	1:	111:39359	
REFERENCE	2:	105:37517	
15	RN	99346-70-0	REGISTRY
REFERENCE	1:	103:224372	
16	RN	99346-63-1	REGISTRY
REFERENCE	1:	103:224372	
17	RN	99346-62-0	REGISTRY
REFERENCE	1:	103:224372	
18	RN	99346-60-8	REGISTRY
REFERENCE	1:	103:224372	
19	RN	61090-75-3	REGISTRY
REFERENCE	1:	86:5484	
20	RN	61090-74-2	REGISTRY
REFERENCE	1:	86:5484	
21	RN	61090-43-5	REGISTRY
REFERENCE	1:	86:5484	
22	RN	50822-08-7	REGISTRY
REFERENCE	1:	79:88264	
23	RN	27677-73-2	REGISTRY
REFERENCE	1:	72:41258	
24	RN	27045-09-6	REGISTRY
REFERENCE	1:	73:3747	
25	RN	27045-07-4	REGISTRY
REFERENCE	1:	73:3747	
26	RN	27045-06-3	REGISTRY
REFERENCE	1:	73:3747	
27	RN	27045-05-2	REGISTRY
REFERENCE	1:	73:3747	
28	RN	27045-04-1	REGISTRY
REFERENCE	1:	73:3747	
29	RN	27045-03-0	REGISTRY

REFERENCE	1:	73:3747	
30	RN	27045-02-9	REGISTRY
REFERENCE	1:	73:3747	
31	RN	27045-01-8	REGISTRY
REFERENCE	1:	73:3747	
32	RN	27045-00-7	REGISTRY
REFERENCE	1:	73:3747	
33	RN	27044-99-1	REGISTRY
REFERENCE	1:	73:3747	
34	RN	19206-24-7	REGISTRY
REFERENCE	1:	71:38935	
35	RN	6758-52-7	REGISTRY
36	RN	6744-43-0	REGISTRY
37	RN	6744-42-9	REGISTRY
38	RN	6739-76-0	REGISTRY
39	RN	6739-75-9	REGISTRY

=> d 1 3 4 6 7 10 ide can

NO ANSWERS DISPLAYED.

THE ANSWER SET WAS CREATED IN FILE 'CA'.

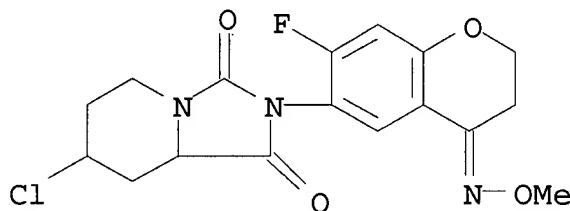
USE THE FILE COMMAND TO CHANGE TO THE CORRECT FILE.

You have entered a file that is not in the current file environment.

Enter "DISPLAY HISTORY" to see a list of the files in the current environment.

=> d 112 1 3 4 6 7 10 ide can

L12 ANSWER 1 OF 39 REGISTRY COPYRIGHT 1996 ACS
RN 174798-58-4 REGISTRY
CN Imidazo[1,5-a]pyridine-1,3(2H,5H)-dione,
7-chloro-2-[7-fluoro-3,4-
dihydro-4-(methoxyimino)-2H-1-benzopyran-6-yl]tetrahydro-
(9CI) (CA
INDEX NAME)
FS 3D CONCORD
MF C17 H17 Cl F N3 O4
SR CA
LC STN Files: CA, CAPLUS

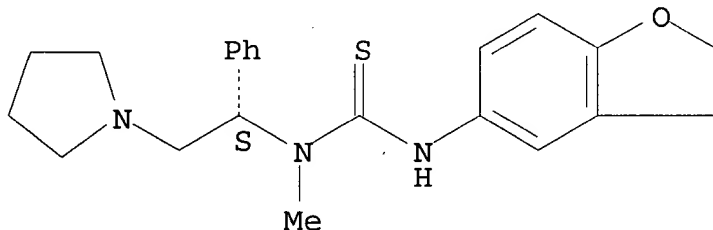


1 REFERENCES IN FILE CA (1967 TO DATE)
 1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 124:261061

L12 ANSWER 3 OF 39 REGISTRY COPYRIGHT 1996 ACS
 RN 149866-40-0 REGISTRY
 CN Thiourea,
 N'-(2,3-dihydro-5-benzofuranyl)-N-methyl-N-[1-phenyl-2-(1-pyrrolidinyl)ethyl]-, monohydrochloride, (S)- (9CI) (CA INDEX NAME)
 FS STEREOSEARCH
 MF C22 H27 N3 O S . Cl H
 SR CA
 LC STN Files: CA, CAPLUS
 DES 1:S

Absolute stereochemistry.



HCl

1 REFERENCES IN FILE CA (1967 TO DATE)
 1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 119:139079

L12 ANSWER 4 OF 39 REGISTRY COPYRIGHT 1996 ACS
 RN 149620-94-0 REGISTRY
 CN Acetic acid,
 [6-[[[(2,3-dihydro-1-methyl-2-oxo-5-phenyl-1H-1,4-benzodiazepin-3-yl)amino]carbonyl]amino]-2,3-dihydro-4H-1-benzopyran-

4-ylidene]-, (E)- (.-+.) - (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 1H-1,4-Benzodiazepine, acetic acid deriv.

FS STEREOSEARCH

MF C28 H24 N4 O5

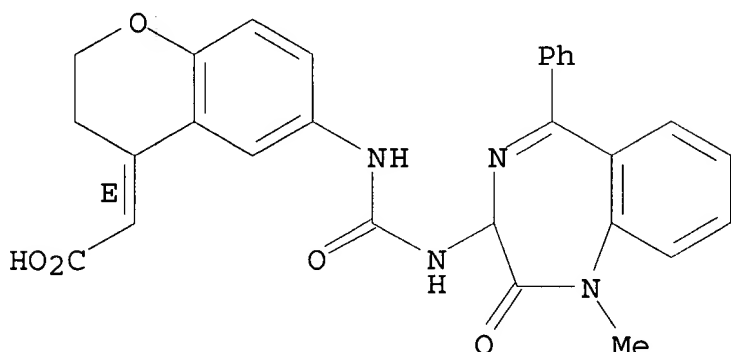
SR CA

LC STN Files: CA, CAPLUS

DES 2:E3:(+-)

Racemate.

Double bond geometry as shown.



1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 119:160327

L12 ANSWER 6 OF 39 REGISTRY COPYRIGHT 1996 ACS

RN 133077-32-4 REGISTRY

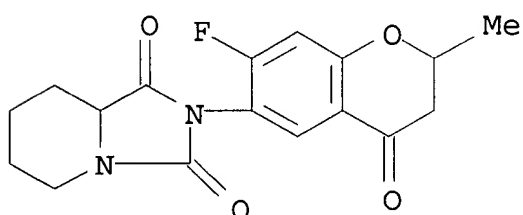
CN Imidazo[1,5-a]pyridine-1,3(2H,5H)-dione,
2-(7-fluoro-3,4-dihydro-2-
methyl-4-oxo-2H-1-benzopyran-6-yl)tetrahydro- (9CI) (CA
INDEX NAME)

FS 3D CONCORD

MF C17 H17 F N2 O4

SR CA

LC STN Files: CA, CAPLUS



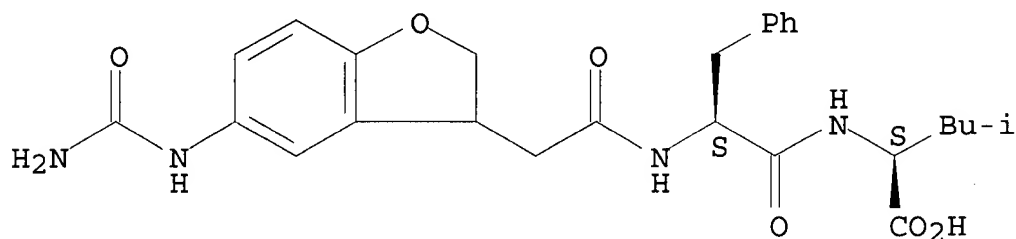
1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 114:164000

L12 ANSWER 7 OF 39 REGISTRY COPYRIGHT 1996 ACS
RN 130105-31-6 REGISTRY
CN L-Leucine, N-[N-[(5-[(aminocarbonyl)amino]-2,3-dihydro-3-
benzofuranyl]acetyl]-L-phenylalanyl]- (9CI) (CA INDEX NAME)
FS STEREOSEARCH
MF C26 H32 N4 O6
SR CA
LC STN Files: CA, CAPLUS
DES 5:L,L

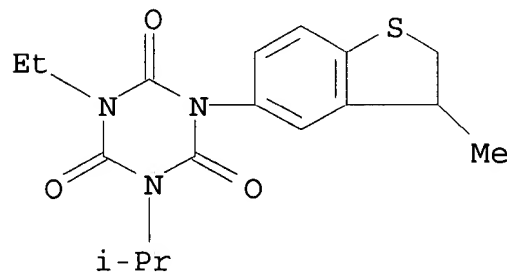
Absolute stereochemistry.



1 REFERENCES IN FILE CA (1967 TO DATE)
1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 113:191969

L12 ANSWER 10 OF 39 REGISTRY COPYRIGHT 1996 ACS
RN 125704-84-9 REGISTRY
CN 1,3,5-Triazine-2,4,6(1H,3H,5H)-trione, 1-(2,3-dihydro-3-
methylbenzo[b]thien-5-yl)-3-ethyl-5-(1-methylethyl)- (9CI)
(CA
INDEX NAME)
FS 3D CONCORD
MF C17 H21 N3 O3 S
SR CA
LC STN Files: CA, CAPLUS, USPATFULL

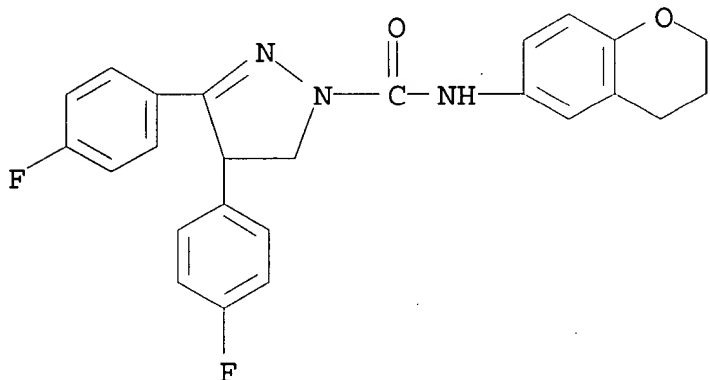


1 REFERENCES IN FILE CA (1967 TO DATE)
1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 112:118858

=> d 112 11 15 19 22 23 24 ide can

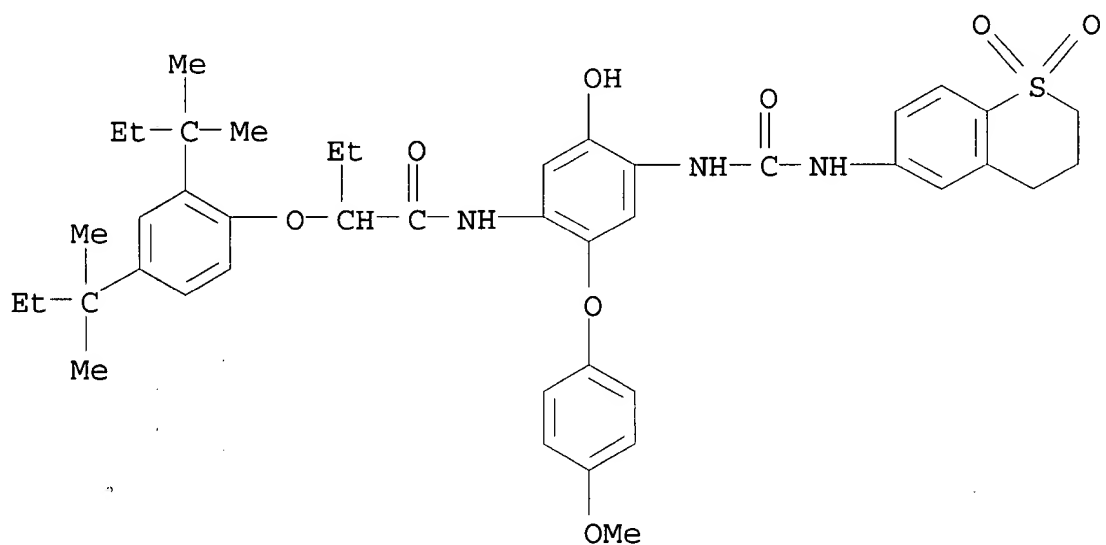
L12 ANSWER 11 OF 39 REGISTRY COPYRIGHT 1996 ACS
RN 121456-04-0 REGISTRY
CN 1H-Pyrazole-1-carboxamide,
N-(3,4-dihydro-2H-1-benzopyran-6-yl)-3,4-
bis(4-fluorophenyl)-4,5-dihydro- (9CI) (CA INDEX NAME)
FS 3D CONCORD
MF C25 H21 F2 N3 O2
SR CA
LC STN Files: CA, CAPLUS, USPATFULL



1 REFERENCES IN FILE CA (1967 TO DATE)
1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 111:39359

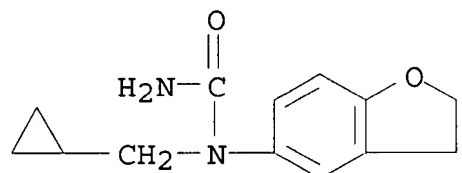
L12 ANSWER 15 OF 39 REGISTRY COPYRIGHT 1996 ACS
RN 99346-70-0 REGISTRY
CN Butanamide,
2-[2,4-bis(1,1-dimethylpropyl)phenoxy]-N-[4-[[[(3,4-
dihydro-2H-1-benzothiopyran-6-yl)amino]carbonyl]amino]-5-hydroxy-
2-
(4-methoxyphenoxy)phenyl]-, S,S-dioxide (9CI) (CA INDEX
NAME)
OTHER CA INDEX NAMES:
CN 2H-1-Benzothiopyran, butanamide deriv.
FS 3D CONCORD
MF C43 H53 N3 O8 S
SR CA
LC STN Files: CA, CAPLUS



1 REFERENCES IN FILE CA (1967 TO DATE)
 1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 103:224372

L12 ANSWER 19 OF 39 REGISTRY COPYRIGHT 1996 ACS
 RN 61090-75-3 REGISTRY
 CN Urea, N-(cyclopropylmethyl)-N-(2,3-dihydro-5-benzofuranyl)-
 (9CI) (CA INDEX NAME)
 FS 3D CONCORD
 MF C13 H16 N2 O2
 LC STN Files: CA, CAPLUS, IFICDB, IFIPAT, IFIUDB, USPATFULL

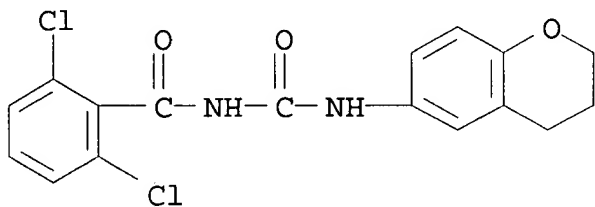


1 REFERENCES IN FILE CA (1967 TO DATE)
 1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 86:5484

L12 ANSWER 22 OF 39 REGISTRY COPYRIGHT 1996 ACS
 RN 50822-08-7 REGISTRY
 CN Benzamide, 2,6-dichloro-N-[[(3,4-dihydro-2H-1-benzopyran-6-yl)amino]carbonyl]- (9CI) (CA INDEX NAME)
 FS 3D CONCORD

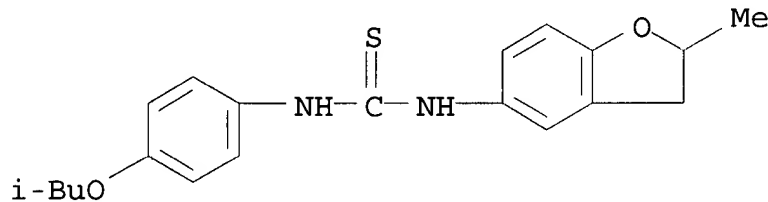
MF C17 H14 Cl2 N2 O3
LC STN Files: CA, CAPLUS, TOXLIT



1 REFERENCES IN FILE CA (1967 TO DATE)
1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 79:88264

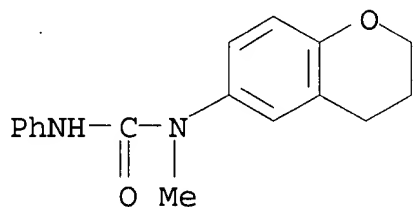
L12 ANSWER 23 OF 39 REGISTRY COPYRIGHT 1996 ACS
RN 27677-73-2 REGISTRY
CN Urea,
1-(2,3-dihydro-2-methyl-5-benzofuranyl)-3-(p-isobutoxyphenyl)-
2-thio- (8CI) (CA INDEX NAME)
FS 3D CONCORD
MF C20 H24 N2 O2 S
LC STN Files: BEILSTEIN*, CA, CAPLUS
(*File contains numerically searchable property data)



1 REFERENCES IN FILE CA (1967 TO DATE)
1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 72:41258

L12 ANSWER 24 OF 39 REGISTRY COPYRIGHT 1996 ACS
RN 27045-09-6 REGISTRY
CN Urea, 1-(6-chromanyl)-1-methyl-3-phenyl- (8CI) (CA INDEX
NAME)
FS 3D CONCORD
MF C17 H18 N2 O2
LC STN Files: BEILSTEIN*, CA, CAPLUS
(*File contains numerically searchable property data)



1 REFERENCES IN FILE CA (1967 TO DATE)
 1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 73:3747

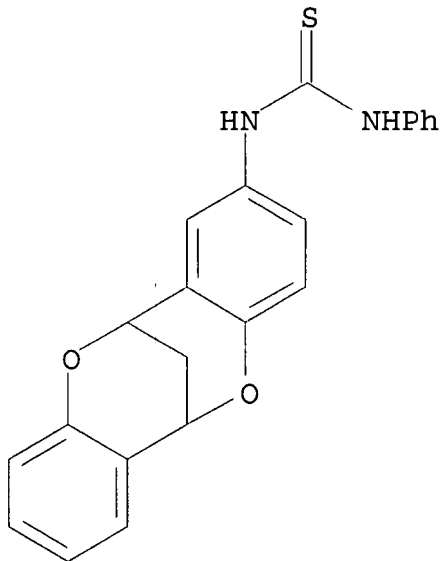
=> d 34-39 ide can
 NO ANSWERS DISPLAYED.

THE ANSWER SET WAS CREATED IN FILE 'CA'.
 USE THE FILE COMMAND TO CHANGE TO THE CORRECT FILE.
 You have entered a file that is not in the current file
 environment.
 Enter "DISPLAY HISTORY" to see a list of the files in the
 current
 environment.

=> d 112 34-39 ide can

L12 ANSWER 34 OF 39 REGISTRY COPYRIGHT 1996 ACS
 RN 19206-24-7 REGISTRY
 CN Urea,
 1-(6,12-methano-6H,12H-dibenzo[b,f][1,5]dioxocin-2-yl)-3-
 phenyl-2-thio-, (.+-.)- (8CI) (CA INDEX NAME)
 FS STEREOSEARCH
 MF C22 H18 N2 O2 S
 LC STN Files: CA, CAPLUS
 DES 3:(+-)

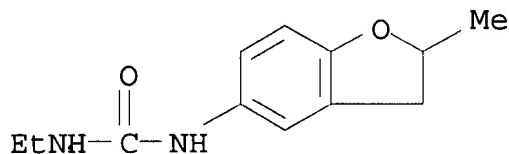
Racemate.



1 REFERENCES IN FILE CA (1967 TO DATE)
 1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

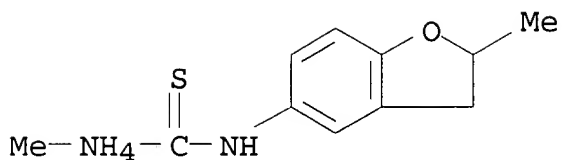
REFERENCE 1: 71:38935

L12 ANSWER 35 OF 39 REGISTRY COPYRIGHT 1996 ACS
 RN 6758-52-7 REGISTRY
 CN Urea, 1-(2,3-dihydro-2-methyl-5-benzofuranyl)-3-ethyl-
 (7CI, 8CI)
 (CA INDEX NAME)
 FS 3D CONCORD
 MF C12 H16 N2 O2
 LC STN Files: CAOLD



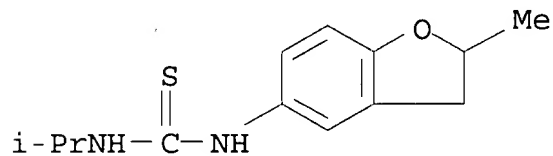
1 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

L12 ANSWER 36 OF 39 REGISTRY COPYRIGHT 1996 ACS
 RN 6744-43-0 REGISTRY
 CN Urea,
 1-(2,3-dihydro-2-methyl-5-benzofuranyl)-3-methyl-2-thio- (7CI,
 8CI) (CA INDEX NAME)
 MF C11 H17 N2 O S
 LC STN Files: CAOLD



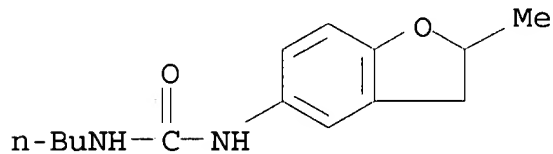
1 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

L12 ANSWER 37 OF 39 REGISTRY COPYRIGHT 1996 ACS
 RN 6744-42-9 REGISTRY
 CN Urea,
 1-(2,3-dihydro-2-methyl-5-benzofuranyl)-3-isopropyl-2-thio-
 (7CI, 8CI) (CA INDEX NAME)
 FS 3D CONCORD
 MF C₁₃ H₁₈ N₂ O S
 LC STN Files: CAOLD



1 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

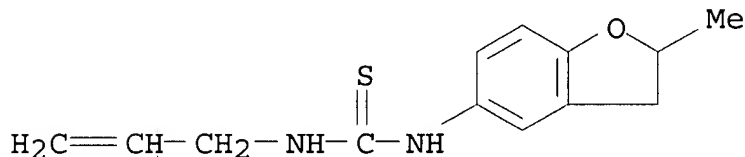
L12 ANSWER 38 OF 39 REGISTRY COPYRIGHT 1996 ACS
 RN 6739-76-0 REGISTRY
 CN Urea, 1-butyl-3-(2,3-dihydro-2-methyl-5-benzofuranyl)-
 (7CI, 8CI)
 (CA INDEX NAME)
 FS 3D CONCORD
 MF C₁₄ H₂₀ N₂ O₂
 LC STN Files: BEILSTEIN*, CAOLD
 (*File contains numerically searchable property data)



1 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

L12 ANSWER 39 OF 39 REGISTRY COPYRIGHT 1996 ACS
 RN 6739-75-9 REGISTRY
 CN Urea,
 1-allyl-3-(2,3-dihydro-2-methyl-5-benzofuranyl)-2-thio- (7CI,

8CI) (CA INDEX NAME)
FS 3D CONCORD
MF C13 H16 N2 O S
LC STN Files: BEILSTEIN*, CAOLD
(*File contains numerically searchable property data)



1 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

=> fil caold
FILE 'CAOLD' ENTERED AT 11:41:41 ON 08 JUL 96
USE IS SUBJECT TO THE TERMS OF YOUR CUSTOMER AGREEMENT
COPYRIGHT (C) 1996 AMERICAN CHEMICAL SOCIETY (ACS)

FILE COVERS 1957-1966
FILE LAST UPDATED: 30 OCT 91 (910803/ED)

To help control your online searching costs, consider using the HCAOLD File when conducting SmartSELECT searches with large numbers of terms.

=> s l12
L14 3 L12

=> d 1-3

L14 ANSWER 1 OF 3 COPYRIGHT 1996 ACS
AN CA65:3835e
DT P
IT 6739-73-7 6739-76-0 6739-77-1

L14 ANSWER 2 OF 3 COPYRIGHT 1996 ACS
AN CA65:3835d
DT P
IT 6739-75-9

L14 ANSWER 3 OF 3 COPYRIGHT 1996 ACS
AN CA65:3835c
DT P
IT 6744-42-9 6744-43-0 6758-52-7

=> fil ca
FILE 'CA' ENTERED AT 11:42:08 ON 08 JUL 96
USE IS SUBJECT TO THE TERMS OF YOUR CUSTOMER AGREEMENT
COPYRIGHT (C) 1996 AMERICAN CHEMICAL SOCIETY (ACS)

FILE COVERS 1967 - 3 Jul 1996 (960703/ED) VOL 125 ISS 2

To help control your online searching costs, consider using the HCA File when using the FSEARCH command or when conducting SmartSELECT searches with large numbers of terms.

Thesauri are now available for the WIPO International Patent Classifications (IPC) editions 1-6 in the /IC1, /IC2, /IC3, /IC4,

/IC5, and /IC (/IC6) fields, respectively. The thesauri in the /IC5 and /IC fields also include the corresponding catchword terms

from the IPC subject headings and subheadings.

*** YOU HAVE NEW MAIL ***

'.CAFILe' IS DEFAULT FORMAT FOR 'CA' FILE

=> d 1 hit

NO ANSWERS DISPLAYED.

THE ANSWER SET WAS CREATED IN FILE 'CAOLD'.

USE THE FILE COMMAND TO CHANGE TO THE CORRECT FILE.

You have entered a file that is not in the current file environment.

Enter "DISPLAY HISTORY" to see a list of the files in the current environment.

=> d 113 1 hit

L13 ANSWER 1 OF 14 CA COPYRIGHT 1996 ACS

IT	174796-99-7P	174797-00-3P	174797-03-6P	174797-04-7P
	174797-05-8P	174797-06-9P	174797-07-0P	174797-08-1P
	174797-09-2P	174797-10-5P	174797-11-6P	174797-12-7P
	174797-13-8P	174797-14-9P	174797-15-0P	174797-16-1P
	174797-17-2P	174797-18-3P	174797-19-4P	174797-20-7P
	174797-21-8P	174797-22-9P	174797-23-0P	174797-24-1P
	174797-25-2P	174797-26-3P	174797-27-4P	174797-28-5P
	174797-29-6P	174797-30-9P	174797-31-0P	174797-32-1P
	174797-33-2P	174797-34-3P	174797-35-4P	174797-36-5P
	174797-37-6P	174797-38-7P	174797-39-8P	174797-40-1P
	174797-41-2P	174797-42-3P	174797-43-4P	174797-44-5P
	174797-45-6P	174797-46-7P	174797-47-8P	174797-48-9P
	174797-49-0P	174797-50-3P	174797-51-4P	174797-52-5P
	174797-53-6P	174797-54-7P	174797-55-8P	174797-56-9P
	174797-57-0P	174797-58-1P	174797-59-2P	174797-60-5P
	174797-61-6P	174797-62-7P	174797-63-8P	174797-64-9P
	174797-65-0P	174797-66-1P	174797-67-2P	174797-68-3P
	174797-69-4P	174797-70-7P	174797-71-8P	174797-72-9P
	174797-73-0P	174797-74-1P	174797-75-2P	174797-76-3P
	174797-77-4P	174797-78-5P	174797-79-6P	174797-80-9P
	174797-81-0P	174797-82-1P	174797-83-2P	174797-84-3P
	174797-85-4P	174797-86-5P	174797-87-6P	174797-88-7P
	174797-89-8P	174797-90-1P	174797-91-2P	174797-92-3P

174797-93-4P 174797-94-5P 174797-95-6P 174797-96-7P
174797-97-8P 174797-98-9P 174797-99-0P 174798-00-6P
174798-01-7P 174798-02-8P 174798-03-9P 174798-04-0P
174798-05-1P 174798-06-2P 174798-07-3P 174798-08-4P
174798-09-5P 174798-10-8P 174798-11-9P 174798-12-0P
174798-13-1P 174798-14-2P 174798-15-3P 174798-16-4P
174798-17-5P 174798-18-6P 174798-19-7P 174798-20-0P
174798-21-1P 174798-22-2P 174798-23-3P 174798-24-4P
174798-25-5P 174798-26-6P 174798-27-7P 174798-28-8P
174798-29-9P 174798-30-2P 174798-31-3P 174798-32-4P
174798-33-5P 174798-34-6P 174798-35-7P 174798-36-8P
174798-37-9P 174798-38-0P 174798-39-1P 174798-40-4P
174798-41-5P 174798-42-6P 174798-43-7P 174798-44-8P
174798-45-9P 174798-46-0P 174798-47-1P 174798-48-2P
174798-49-3P 174798-50-6P 174798-51-7P 174798-52-8P
174798-53-9P 174798-54-0P 174798-55-1P 174798-56-2P

174798-57-3P 174798-58-4P 174798-59-5P

174798-60-8P

RL: AGR (Agricultural use); SPN (Synthetic preparation);

BIOL

(Biological study); PREP (Preparation); USES (Uses)
(prepn. of

2-phenyl-7-chloroperhydroimidazo[1,5-a]pyridine
herbicides for controlling undesired weeds)

=> d 1-14 cbib,abs

NO ANSWERS DISPLAYED.

THE ANSWER SET WAS CREATED IN FILE 'CAOLD'.

USE THE FILE COMMAND TO CHANGE TO THE CORRECT FILE.

You have entered a file that is not in the current file environment.

Enter "DISPLAY HISTORY" to see a list of the files in the current environment.

=> d 113 cbib abs

L13 ANSWER 1 OF 14 CA COPYRIGHT 1996 ACS

124:261061 Preparation of 2-phenyl-7-chloroperhydroimidazo[1,5-a]pyridine herbicides for controlling undesired weeds.

Seckinger,

Karl; Mohanty, Sasank Sekhar; Milzner, Karlheinz; Kuhnhen,

Fred

(Sandoz Ltd., Switz.; Sandoz-Patent-GmbH; Sandoz-Erfindungen Verwaltungsgesellschaft m.b.H.). Eur. Pat. Appl. EP 688773

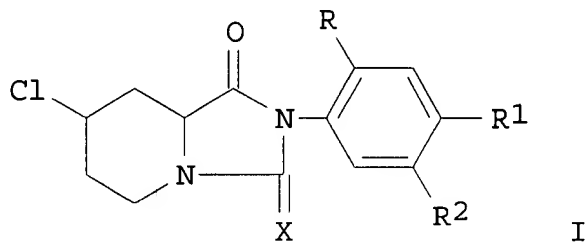
A1

951227, 24 pp. DESIGNATED STATES: R: AT, BE, CH, DE, DK, ES, FR,

GB, GR, IE, IT, LI, LU, NL, PT, SE. (English). CODEN:
EPXXDW.

APPLICATION: EP 95-810410 950620. PRIORITY: GB 94-12603
940623.

GI



AB The title compds. (I; X = O, S; R = H, Cl, F; R1 = F, Cl, Br, CN,

Me; R2 = halogen, C 1-6 alkyl, C1-6 alkoxy, C1-6 alkylcarbonyloxy, C3-6 cycloalkoxy, C3-6 alkynyloxy, C3-6 alkenyloxy, CO2H, etc.),

useful as herbicides for the control of undesired weeds, are prep'd.

Thus, 4-chloro-2-piperidinecarboxylic acid Me ester hydrochloride

was reacted with the isocyanate of Me 2-chloro-4-fluoro-5-aminocinnamate, producing herbicidal Me 2-chloro-4-fluoro-5-(7-

chloroperhydroimidazo[1,5-a]pyridine-1,3-dione-2-yl)cinnamate, m.p.

162-163.degree..

=> d l13 2-14 cbib abs

L13 ANSWER 2 OF 14 CA COPYRIGHT 1996 ACS

119:160327 Preparation of 3-ureidobenzodiazepinones useful as CCK or

gastrin antagonists. Capet, Marc; Cotrel, Claude; Dubroeucq, Marie

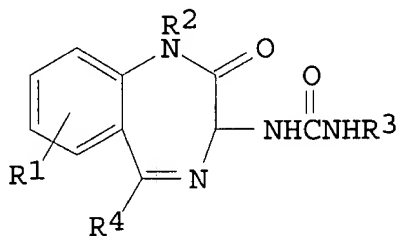
Christine; Guyon, Claude; Martin, Jean Paul (Rhone-Poulenc Rorer SA,

Fr.). Eur. Pat. Appl. EP 538099 A1 930421, 31 pp.

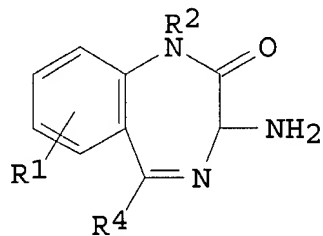
DESIGNATED

STATES: R: PT. (French). CODEN: EPXXDW. APPLICATION: EP 92-402741 921008. PRIORITY: FR 91-12481 911010.

GI



I



II

AB Title compds. I [R1 = H, halo, alkyl, alkoxy, alkylthio, NO₂, OH,
-CN; R2 = alkyl, CHR₅COR₆ (R₅ = H, alkyl, alkoxy carbonyl, various

(un)substituted Ph, R₆ = alkoxy, various (un)substituted cycloalkyloxy, cycloalkylalkyloxy, various substituted N derivs.,

cyclic and acyclic); R₃ = Ph substituted by one or more ZSO₃H (Z =

alkylene), ZPO₃H₂, CH:NOH, CHNOZCO₂X, SOZCO₂X, SZCO₂X, SO₂ZCO₂X,

CH:CHCO₂X, ZCONHOH, C(:NOH)CO₂X, ZN(OH)CO₂Z, ZSO₂H,

CH:CHSO₃H,
C(CO₂X):NOZCO₂X, tetrazolylalkyl, etc.] are prep'd. as CCK or gastrin

antagonists (no data) by condensation of a carbonic acid deriv. and

amine R₃NH₂ with an aminodihydrobenzodiazepinone II.

L13 ANSWER 3 OF 14 CA COPYRIGHT 1996 ACS

119:139079 Preparation of (pyrrolidinoethyl)urea derivatives as analgesics. Takeuchi, Makoto; Takayama, Kazuhisa; Onda, Kenichi;

Motoie, Hiroyuki; Isomura, Yasuo (Yamanouchi Pharmaceutical Co.,

Ltd., Japan). PCT Int. Appl. WO 9303011 A1 930218, 93 pp.

DESIGNATED STATES: W: AT, AU, BB, BG, BR, CA, CH, CS, DE, DK, ES,

FI, GB, HU, JP, KR, LK, LU, MG, MN, MW, NL, NO, PL, RO, RU, SD, SE,

US; RW: AT, BE, BF, BJ, CF, CG, CH, CI, CM, DE, DK, ES, FR, GA, GB,

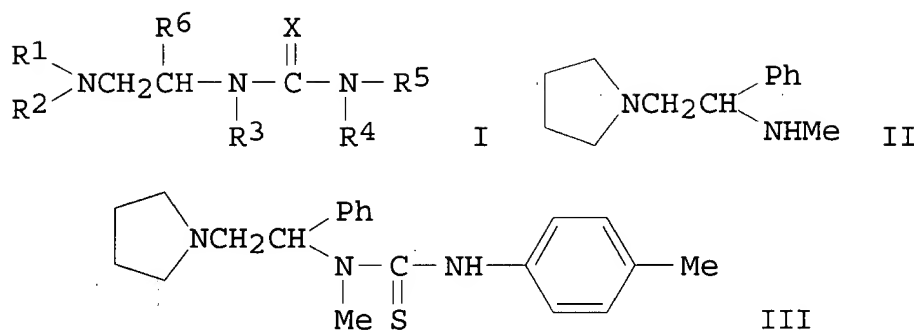
GR, IE, IT, LU, MC, ML, MR, NL, SE, SN, TD, TG.

(Japanese). CODEN:

PIXXD2. APPLICATION: WO 92-JP993 920804. PRIORITY: JP
91-223280

910808; JP 91-309952 911029.

GI



AB The title compds. [I; R₁, R₂ = alkyl, alkenyl, alkynyl, cycloalkyl,

R₁R₂N pyrrolidino; R₃, R₄ = H, alkyl, alkenyl, alkynyl, cycloalkyl;

R₃R₄ = alkylene, alkenylene, etc.; R₅ = (substituted) carbocyclic,

condensed heterocyclyl contg. 1 or 2 O and/or S atoms; R₆ = (substituted) Ph; X = O, S] are prep'd. A mixt. of 4-MeC₆H₄NCS and

pyrrolidine deriv. (S)-II in ClCH₂CH₂Cl was stirred at room temp. to

give thiourea (S)-III, which was treated with 4N HCl in EtOAc to

give (S)-III.HCl. III.HCl showed EO50 of 0.54 mg/kg s.c. in mice in

the tail pinch test. Tablet, capsule, injection formulations were given.

L13 ANSWER 4 OF 14 CA COPYRIGHT 1996 ACS

114:164000 Preparation of N-aryl imides as herbicides.

Kunisch, Franz;

Arlt, Dieter; Santel, Hans Joachim; Luerssen, Klaus; Schmidt, Robert

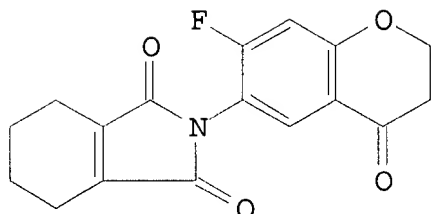
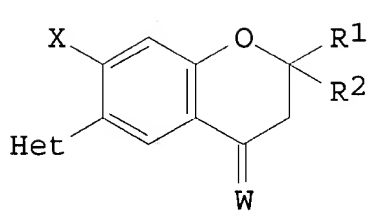
R. (Bayer A.-G., Fed. Rep. Ger.). Eur. Pat. Appl. EP 400403 A2

901205, 37 pp. DESIGNATED STATES: R: BE, CH, DE, FR, GB, IT, LI,

NL. (German). CODEN: EPXXDW. APPLICATION: EP 90-109300 900517.

PRIORITY: DE 89-3917515 890530.

GI



AB N-Aryl imide derivs. I [R1,R2 = H, alkyl; Het = (substituted)
tetrahydropthalimido, maleimido, and other cyclic N- and O-contg.
imides; X = H, halo; W = O, NOR3; R3 = H, (substituted)
alkyl,
alkenyl, alkynyl or cycloalkyl], useful as herbicides (no
data),
were prep'd. For example, a mixt. of
3,4,5,6-tetrahydronphthalic
anhydride and 6-amino-7-fluorochroman-4-one (prepn. given)
in HOAc
was refluxed 3 h to give 42% imide II. Various I show
better
activity as post-emergent herbicides when compared to a
known
herbicide.

L13 ANSWER 5 OF 14 CA COPYRIGHT 1996 ACS
113:191969 Renin inhibitory peptides containing
(4S)-amino-5-cyclohexyl-
(3S)-hydroxypentanoic acid. Smith, Stephen Allan; Ham,
Peter; Nash,
David John (Beecham Group PLC, UK). Eur. Pat. Appl. EP
350163 A2
900110, 91 pp. DESIGNATED STATES: R: AT, BE, CH, DE, ES,
FR, GB,
GR, IT, LI, LU, NL, SE. (English). CODEN: EPXXDW.

APPLICATION: EP
89-305691 890606. PRIORITY: GB 88-13671 880609; GB
88-29065 881213;
GB 89-6262 890318.
GI For diagram(s), see printed CA Issue.
AB The title peptides [I; Z1Z2Z3 = atoms to complete a
5-membered
nonarom. heterocyclic ring; E = absent, $(CH_2)_n$, $CH(CH_2)_{n-1}$;
 $n = 1-4$;
A = CONH, NHCO, CO₂, CH₂, S(O)_r; r, p = 0-2; q = 0,1; R1 =
(un)substituted (hetero)aryl methyl; R2 = CHR₈R₉; R₈ = H, Me
and R₉ =
C₁₋₆ alkyl, C₃₋₈ cycloalkyl, (un)substituted (hetero)aryl;
R₉ = NH₂,

C2-7 alkanoylamino, 2-oxopyrrolidinyl, etc.; R3 = alkyl, cycloalkylmethyl; R4 = (cyclo)alkyl; R5 = H, alkyl; or R5 = OH when

A = CH₂; R6, R7 = H, substituent], useful for the treatment of

hypertension, are prep'd. Thus, N-(2,3-dihydrobenzofuran-2-carbonyl)-(S)-phenylalanyl-(S)-leucine was condensed with (4S)-amino-5-cyclohexyl-(3S)-hydroxypentanoic acid isobutylamide (ACHPAA) in the presence of hydroxybenzotriazole and DCC in THF at room temp.

overnight to give Q-Phe-Leu-ACHPAA (II; Q = 2,3-dihydrobenzofuran-2-carbonyl). II [Q = (6-aminomethyl-2,3-dihydro-1,1-dioxobenzothiophen-3-ylacetyl] in vitro inhibited human renin with

an IC₅₀ of 0.8 times. 10-8M. A total of 75 I were prep'd.

L13 ANSWER 6 OF 14 CA COPYRIGHT 1996 ACS

112:118858 Trisubstituted 1,3,5-triazine-2,4,6-triones as agrochemical

fungicides. Adler, Alfons; Widdig, Arno; Kuehle, Engelbert; Fuehrer, Wolfgang; Hagemann, Hermann; Haenssler, Gerd (Bayer A.-G., Fed. Rep. Ger.). Eur. Pat. Appl. EP 334135 A2 890927, 32 pp.

DESIGNATED STATES: R: BE, CH, DE, FR, GB, IT, LI, NL.
(German).

CODEN: EPXXDW. APPLICATION: EP 89-104338 890311.
PRIORITY: DE
88-3810080 880325.

GI For diagram(s), see printed CA Issue.

AB The title compds [I; R₁ = (substituted) aliph., arom., or cycloaliphatic residue; R₂ = (substituted) aliph. residue; R₃ = (substituted) benzoheterocyclyl], useful as pesticides, were prep'd.

Thus, MeI was added to a mixt. of 1-(2,2-dimethylpropyl)-3-[6-

(2,2,3,3-tetrafluoro-2,3-dihydro-1,4-benzodioxinyl)]-1,3,5-triazine-

2,4,6-trione. The mixt. was refluxed 5 h to give 88% triazinetrione

II. Several I as 0.025% sprays gave 90-100% control of Pyricularia oryzae on rice.

L13 ANSWER 7 OF 14 CA COPYRIGHT 1996 ACS

111:39359 Insecticidal pyrazoline-1-carboxamides, compositions containing them, and their use. Duggan, Angelina J. (FMC Corp.,

USA). U.S. US 4767779 A 880830, 21 pp. Cont.-in-part of U.S. Ser.

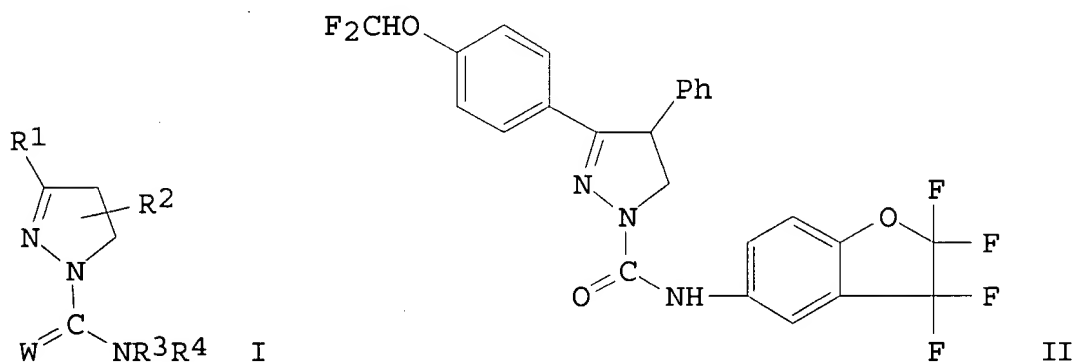
No. 779,721, abandoned. (English). CODEN: USXXAM.

APPLICATION: US

86-849658 860409. PRIORITY: US 84-664674 841025; US 85-709626

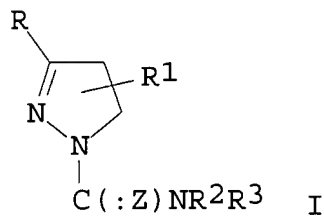
850308; US 85-779721 850924.

GI



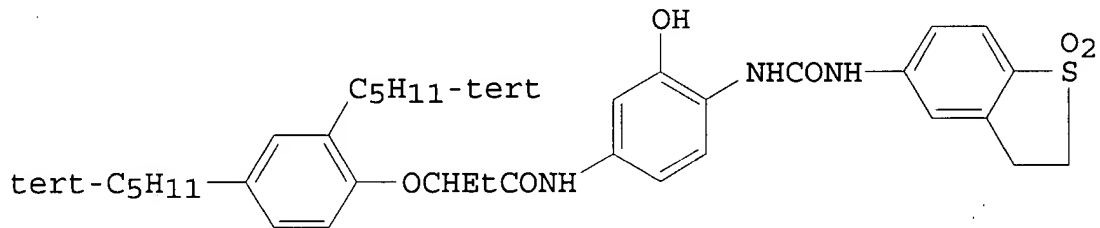
AB The title compds. [I; R1, R2 = (un)substituted Ph optionally fused by O-contg. satd. ring; R3 = (un)substituted C₆H₄OPh, C₆H₄SPh, Ph fused by O-contg. satd. ring, indanyl; R4 = H, alkyl; W = O, S] are prep'd. as insecticides. Etherification of 2,4-Cl(O₂N)C₆H₃OH with BrCF₂CF₂Br in DMF in the presence of K₂CO₃ and PrSH at 50/.degree. gave 2,4-Cl(O₂N)C₆H₃OCF₂CF₂Br, which was cyclized by powd. Cu and 2,2'-bipyridyl in DMSO at 190-195.degree. to give 2,3-dihydro-2,2,3,3-tetrafluoro-5-nitrobenzofuran. This underwent hydrogenation over PtO₂ to give the 5-amino compd., which was treated with COCl₂ in refluxing PhMe to give the isocyanate. Reaction of the latter with 3-(4-difluoromethoxyphenyl)-4-phenylpyrazoline in Et₂O contg. Et₃N catalyst gave (dihydrotetrafluorobenzofuranyl)(difluoromethoxyphenyl)phenylpyrazolinecarboxamide II. As an 8-ppm foliar spray on pinto bean plants prior to infestation, II was 100% lethal to Spodoptera eridania, S. exigua, and Epilachna varivestis, and 95% lethal to Trichoplusia ni.

L13 ANSWER 8 OF 14 CA COPYRIGHT 1996 ACS
105:37517 Pyrazoline insecticides. Duggan, Angeline Joy (FMC Corp., USA). Ger. Offen. DE 3537884 A1 860430, 61 pp. (German).
CODEN: GWXXBX. APPLICATION: DE 85-3537884 851024. PRIORITY: US 84-664674
841025; US 85-709626 850308; US 85-779721 850924.
GI



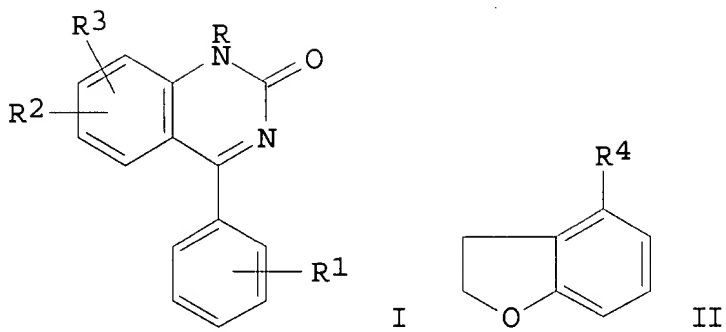
AB The pyrazolines I [R = (un)substituted Ph, etc.; R1 = (un)substituted Ph, 1,4-benzodioxan-6-yl, 1,3-benzodioxol-5-yl, etc.; R2 = H, alkyl; R3 = (un)substituted PhOC6H4, (un)substituted 1,3-benzodioxol-5-yl, benzofuran-5-yl, etc.; Z = O, S] are prepd. as insecticides. Thus, 4-FC6H4NO2 was condensed with 4-ClC6H4OH in K2CO3-contg. DMSO at 70.degree. to give 4-(4-ClC6H4O)C6H4NO2, which was hydrogenated into 4-(4-ClC6H4O)C6H4NH2 on PtO2. The amine was reacted with ClCO2CCl3 in PhMe and the product treated with 3-(4-chlorophenyl)-4-phenylpyrazoline to give I [R = 4-ClC6H4; R1 = 4-Ph, R2 = H; R3 = 4-(4-ClC6H4O)C6H4; Z = O] (II). II (500 ppm) controlled Spodoptera ridania and Epilachna varivestis, on bean leaves, in the lab.

L13 ANSWER 9 OF 14 CA COPYRIGHT 1996 ACS
103:224372 Silver halide photographic material. (Konishiroku Photo Industry Co., Ltd., Japan). Jpn. Kokai Tokkyo Koho JP 60108846 A2
850614 Showa, 10 pp. (Japanese). CODEN: JKXXAF.
APPLICATION: JP
83-218222 831118.
GI



AB A Ag halide photog. material has .gtoreq.1 emulsion layer contg. a phenolic cyan coupler having an arylureido group having a Ph ring to which a heterocyclic ring is condensed through -S- or -SO₂- (the -S- or -SO₂- is directly linked with the phenol ring) at the 2-position, a H or a coupling-off group at the 4-position, and an acylamino group at the 5-position of the phenol ring. By reacting with an oxidized developing agent, it forms a cyan dye which has a sharp spectral absorption in the red region with a low level of unwanted green absorption. The dye-forming activity is also insensitive to benzyl alc. concn. in a developer or to the exhaustion of processing solns. Thus, a Ag(Br,I) emulsion (AgI 5 mol%) contg. I had a good developability upon development by a typical color neg. process, even when a fairly exhausted bleach soln. was used, and formed a cyan dye image with excellent spectral absorption.

L13 ANSWER 10 OF 14 CA COPYRIGHT 1996 ACS
 86:5484 Tricyclic furoquinazolinones. Cooke, George A.;
 Houlihan,
 William J. (Sandoz-Wander, Inc., USA). U.S. US 3963717
 760615, 11 pp. (English). CODEN: USXXAM. APPLICATION: US 75-556574
 750310.
 GI



AB Antiinflammatory and analgesic (no data) furoquinazolinones

I ($R =$
CHMe₂, cyclopropylmethyl, cyclopentylmethyl, CMe₃,
CH₂CMe:CH₂, Et;
 $R_1 = H, 4\text{-F}, 4\text{-CF}_3, 3\text{-OMe}; R_2R_3 = 7,8\text{-OCH}_2\text{CH}_2, 6,7\text{-OCH}_2\text{CH}_2,$
 $5,6\text{-CH}_2\text{CH}_2\text{O}, 6,7\text{-CH}_2\text{CH}_2\text{O}, 5,6\text{-OCH}_2\text{CH}_2, 7,8\text{-CH}_2\text{CH}_2\text{O}$) (38
compds.)

were prep'd. Thus the benzofuranamine II ($R_4 = NH_2$) was treated with

Me₂CHI, II ($R_4 = NHCHMe_2$) treated with NaNCO, II [$R_4 = N(CHMe_2)CONH_2$] condensed with PhCHO and oxidized with KMnO₄ to give

I ($R = CHMe_2, R_1 = H, R_2R_3 = 7,8\text{-OCH}_2\text{CH}_2$).

L13 ANSWER 11 OF 14 CA COPYRIGHT 1996 ACS

79:88264 Synthesis and laboratory evaluation of
1-(2,6-disubstituted

benzoyl)-3-phenylureas, a new class of insecticides. I.

1-(2,5-Dichlorobenzoyl)-3-phenylureas. Wellinga, Kobus;

Mulder,

Rudolf; Van Daalen, Jan J. (Res. Lab., Philips-Duphar B.V.,
Weesp,

Neth.). J. Agr. Food Chem., 21(3), 348-54 (English) 1973.

CODEN:

JAFCAU.

AB Addnl. data considered in abstracting and indexing are available

from a source cited in the original document. Out a large no. of

1-(2,6-dichlorobenzoyl)-3-phenylureas I ($R =$ mono-, di- or trihalo,

alkyl, chloroalkyl, or aryl, $R_1 = H$, alkyl, haloalkyl, or alkenyl,

$R_2 = H, Me, OMe, PhCH_2$ or OH) sensitized and tested against Aedes

aegypti, Pieris brassicae and Leptinotarsa decemlineata,

1-(2,6-dichlorobenzoyl)-3-(4-chlorophenyl)urea (I, $R = 4\text{-Cl}, R_1 = R_2$

= H) [35409-97-3] was the most active. In many cases, the

activities against the 3 test insects differed, *M. decemlineata* being usually the least sensitive. When R was dihalo, the lowest activities were shown in position 2,6. High activity was shown for R = alkyl. I showed the highest activities when R1 and R2 were H. Very poor activity was shown when R was an electron-attracting group. I acted by disturbing the cuticle deposition, resulting in abortive molt.

L13 ANSWER 12 OF 14 CA COPYRIGHT 1996 ACS
73:3747 Substituted chroman-6-ylureas and thioureas. Lettieri, G.; Brancaccio, Giovanni; Larizza, Angelo; Viterbo, Rene (Res. Lab., Richardson-Merrell S.p.A., Naples, Italy). J. Med. Chem., 13(3), 584-5 (English) 1970. CODEN: JMCMAR.
AB I (R = H, Me, or Cl; R1 = H or Me; R2 = Ph, 4-ClC₆H₄, Pr, 3-(O₂N)C₆H₄, 2-MeOC₆H₄, or 4-EtOC₆H₄; X = O or S) are prepd. from chromanylaminates and isocyanates or isothiocyanates.

L13 ANSWER 13 OF 14 CA COPYRIGHT 1996 ACS
72:41258 Tuberculostatic 1,3-diarylthioureas. I. Winkelmann, Erhardt; Wagner, Wolf H.; Hilmer, Hans (Farbwerke Hoechst A.-G., Frankfurt/M.-Hoechst, Ger.). Arzneim.-Forsch., 19(4), 543-58 (German) 1969. CODEN: ARZNAD.
AB One hundred eighty different Ph substituted thioureas (R₁NHCSNH r₂) were tested for tuberculostatic activity in vitro and in the mouse. The tables presented indicate that p-BuOC₆H₄NHCSNHC₆H₄OBu-m (I) had the greatest activity in vitro (0.1-0.2 .mu.g/ml) while in vivo I was most active at a dosage of 250 mg/kg body wt. when given orally.

L13 ANSWER 14 OF 14 CA COPYRIGHT 1996 ACS
71:38935 6H,12H-6,12-methanodibenzo[b,f] [1,5]dioxocins from the reactions of o-coumaric acids and salicylaldehydes. Hennis, Henry E.; Wang, Chun-Shan (Benzene Res. Lab., Dow Chem. Co., Midland, Mich., USA). J. Org. Chem., 34(6), 1907-11 (English) 1969. CODEN:

JOCEAH.

AB 6H,12H-6,12-Methanodibenzo[b,f][1,5]dioxocin (I) was prep'd. from the

reaction of o-vinylphenol or o-coumaric acid (II) and salicylaldehyde (III) in 4.6% yield. 2-Methyl-, 2-bromo-(IV), and

2-nitro-6H,12H-6,12-methanodibenzo[b,f][1,5]-dioxocins (V) were

synthesized by the reactions of 2-hydroxy-5-methylcinnamic acid with

III and II with 5-bromo- and 5-nitro-salicylaldehyde, resp.

The

reactions of 2-hydroxy-1-naphthaldehyde gave heterocyclics contg. a

naphthalene ring. I was brominated to the 2,8-dibromo deriv. (VI).

Both IV and VI were converted into the nitriles by reaction with

CuCN. Neither IV nor VI could be converted into Grignard reagents,

but were readily metallated with BuLi. The organometallics were

carbonated to yield carboxylic acids. The heterocyclic ring system

of I was cleaved by hydrogenolysis to 2,2'-trimethylenediphenol.

(.+-.)-2-Amino-6H,12H-6,12-methanodibenzo[b,f][1,5]dioxocin (.+-.-VII), prep'd. by the catalytic hydrogenation of V, was resolved

via the tartrate salts to yield the optical isomers, [.alpha.]_{25D}

389.0 and -393.3.degree.. The more abundant (+)-VII was reduced via

diazotization to (+)-I, [.alpha.]_{25D} 266.7.degree..

=> fil reg

FILE 'REGISTRY' ENTERED AT 11:46:34 ON 08 JUL 96

USE IS SUBJECT TO THE TERMS OF YOUR CUSTOMER AGREEMENT

COPYRIGHT (C) 1996 American Chemical Society (ACS)

STRUCTURE FILE UPDATES: 6 JUL 96 HIGHEST RN 177929-67-8

DICTIONARY FILE UPDATES: 6 JUL 96 HIGHEST RN 178150-05-5

TSCA INFORMATION NOW CURRENT THROUGH DECEMBER 1995

Please note that search-term pricing does apply when conducting SmartSELECT searches.

'REGISTRY' IS DEFAULT FORMAT FOR 'REGISTRY' FILE

=> d 23

NO ANSWERS DISPLAYED.

THE ANSWER SET WAS CREATED IN FILE 'CAOLD'.

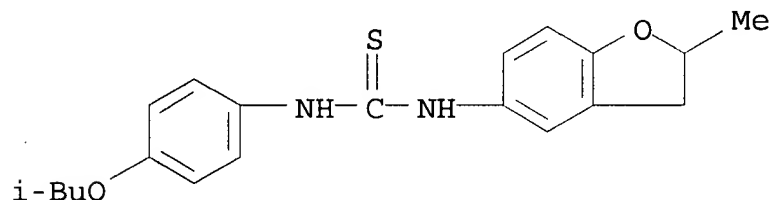
USE THE FILE COMMAND TO CHANGE TO THE CORRECT FILE.

You have entered a file that is not in the current file environment.

Enter "DISPLAY HISTORY" to see a list of the files in the current environment.

=> d 112 23

L12 ANSWER 23 OF 39 REGISTRY COPYRIGHT 1996 ACS
RN 27677-73-2 REGISTRY
IN Urea,
1-(2,3-dihydro-2-methyl-5-benzofuranyl)-3-(p-isobutoxyphenyl)-
2-thio- (8CI)
MF C20 H24 N2 O2 S
LC STN Files: BEILSTEIN*, CA, CAPLUS
(*File contains numerically searchable property data)



1 REFERENCES IN FILE CA (1967 TO DATE)
1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1

AN 72:41258 CA
SO Arzneim.-Forsch. (1969), 19(4), 543-58
CODEN: ARZNAD
PY 1969